Conformational Transitions of Subunit ε in ATP Synthase from Thermophilic Bacillus PS3

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ABSTRACT Subunit ϵ of bacterial and chloroplast F_0F_1 -ATP synthase is responsible for inhibition of ATPase activity. In Bacillus PS3 enzyme, subunit ε can adopt two conformations. In the "extended", inhibitory conformation, its two C-terminal α -helices are stretched along subunit γ . In the "contracted", noninhibitory conformation, these helices form a hairpin. The transition of subunit ε from an extended to a contracted state was studied in ATP synthase incorporated in Bacillus PS3 membranes at 59°C. Fluorescence energy resonance transfer between fluorophores introduced in the C-terminus of subunit ε and in the N-terminus of subunit γ was used to follow the conformational transition in real time. It was found that ATP induced the conformational transition from the extended to the contracted state (half-maximum transition extent at 140 μ M ATP). ADP could neither prevent nor reverse the ATP-induced conformational change, but it did slow it down. Acid residues in the DELSEED region of subunit β were found to stabilize the extended conformation of ε . Binding of ATP directly to ε was not essential for the ATPinduced conformational change. The ATP concentration necessary for the half-maximal transition (140 μ M) suggests that subunit ε probably adopts the extended state and strongly inhibits ATP hydrolysis only when the intracellular ATP level drops significantly below the normal value.

INTRODUCTION

 H^+ -F_OF₁-ATP synthase is found in bacterial plasma membranes, thylakoid membranes of chloroplasts, and inner mitochondrial membranes ([1,2](#page-7-0)). The enzyme uses the energy of the transmembrane electrochemical proton potential difference $(\Delta \tilde{\mu}_{H^+})$ to synthesize ATP from ADP and P_i. The simplest forms of bacterial F_0F_1 , including *Bacillus* PS3 enzyme, have eight types of subunits. Five subunits (in stoichiometry $\alpha_3\beta_3\gamma_1\delta_1\epsilon_1$) comprise hydrophilic F₁, which bears nucleotide-binding sites. Three subunits (in stoichiometry $a_1b_2c_{10}$ comprise membrane-embedded F_O, which is responsible for proton translocation. Three $\alpha\beta$ pairs form a hexamer, and elongated subunit γ fills its central cavity ([3–5\)](#page-7-0). Subunits c form a ring-shaped oligomer [\(6–8](#page-7-0)) that binds to the $\gamma \varepsilon$ -complex [\(6,9\)](#page-7-0).

During catalysis, proton translocation powers the rotation of the c-oligomer (and the $\gamma \varepsilon$ -complex bound to it) relative to the $ab_2\alpha_3\beta_3\delta_1$ -complex [\(10–13](#page-7-0)). Rotation of the $\gamma\epsilon$ -complex inside the $\alpha_3\beta_3$ -hexamer induces cyclic conformational changes in catalytic sites that result in substrate binding and product release [\(14–17](#page-7-0)).

When the $\Delta \tilde{\mu}_{H^+}$ is low, the enzyme operates as an ATPdriven proton pump. This activity reversal can be important for such functions as maintaining the $\Delta\widetilde{\mu}_{H^+}$ necessary for ion transport and flagella motility in bacteria or pumping out H^+ in acidophiles. However, uncontrolled ATPase activity can quickly exhaust the intracellular ATP pool. Several regulatory features prevent this by selectively inhibiting ATP

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hydrolysis, but not synthesis ([2](#page-7-0)), converting the protein from a mere catalyst that speeds up both forward and reverse reactions into a unidirectional molecular machine.

One such feature found in bacterial and chloroplast F_0F_1 -ATP synthase is the inhibition of ATPase activity by subunit ε [\(18–20](#page-8-0)). Subunit ε is a small, 15 kDa protein consisting of two domains: the N-terminal β -sandwich and the C-terminal pair of α -helices. Only the latter domain is responsible for the inhibition [\(21–28](#page-8-0)). Numerous studies have revealed that subunit ϵ can undergo large conformational changes ([25,29–35](#page-8-0)). Some details of these conformational changes were elucidated by x-ray and NMR studies demonstrating that the C-terminal domain of ε can adopt at least two conformations: a hairpin-like contracted state (the ''down state''; [Fig. 1](#page-1-0) B, state III) [\(6,33,36–38](#page-7-0)) and an extended state with the two α -helices stretched along subunit γ toward the $\alpha_3\beta_3$ -hexagon (the "up state"; [Fig. 1](#page-1-0) B, state I) [\(39,40\)](#page-8-0). In later studies, these two different states were trapped by crosslinking in F_0F_1 from *Escherichia coli* ([41\)](#page-8-0) and *Bacillus* PS3 ([42\)](#page-8-0). It was also found that fixing subunit ε in the contracted hairpin-like state abolishes the inhibitory effect ([41,42](#page-8-0)).

Experiments on Bacillus PS3 enzyme have been particularly fruitful in elucidating the inhibitory function of subunit ε . It was found that in the presence of ATP, subunit ε adopts a noninhibitory contracted conformation, whereas in the presence of ADP, the inhibitory extended state is observed. As a possible explanation, it was suggested that the ATP/ADP ratio determines the conformation of subunit ε ([42\)](#page-8-0).

It was also suggested that electrostatic interactions between the positively charged residues in the C-terminal domain of ε and the negatively charged residues in the

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FIGURE 1 (A) Stereo cartoon representation of subunit ε (dark gray) in an extended conformation. Partly shown are subunit γ (white) and subunit β proximal to the C-terminal domain of ε . Fluorescent labels are shown as large colored spheres (magenta: Cy3 on the N-terminus of γ ; blue: Cy5 on the C-terminus of ε). Residues mutated to reduce the affinity of subunit ε to ATP are shown as small orange spheres, and the β DELSDED region is shown in green. The figure was generated from Protein Data Bank entry 1JNV of the E. coli $\alpha_3\beta_3\gamma\epsilon$ complex ([40\)](#page-8-0) with VMD software ([53\)](#page-8-0). (B) Cartoon scheme of subunit ε conformational transitions in Bacillus PS3 F_0F_1 ATP synthase. Subunit γ is shown in white, subunit ε in dark gray, and $\alpha_3\beta_3\delta$ -complex and F_O in light gray. Positions of introduced Cy3 on subunit γ and Cy5 on subunit ε are indicated by a magenta and blue star, respectively. See details at the end of the Discussion.

DELSDED region of subunit β (green in Fig. 1 A) are involved in propagating the inhibitory effect of subunit ε ([43\)](#page-8-0). However, it remained unclear whether the electrostatic interactions with β DELSDED play any role in the conformational changes of ε .

A peculiar feature of *Bacillus* PS3 F_0F_1 is that ATP can bind directly to subunit ε [\(44,45\)](#page-8-0). The recently published structure of the ε -ATP complex confirms that ATP binds when subunit ϵ is in the hairpin noninhibitory conformation ([46\)](#page-8-0). Mutagenesis experiments revealed that weaker binding of ATP to isolated subunit ϵ correlated with greater inhibition of $\alpha_3\beta_3\gamma_5$ -complex ATPase activity, suggesting a regulatory role for ATP binding to ε in vivo [\(47](#page-8-0)).

To our knowledge, the detailed mechanism and kinetics of subunit ϵ conformational transitions in the whole F_0F_1 from Bacillus PS3 have not yet been studied. However, some results have been obtained on the isolated F_1 portion. Iino et al. ([48\)](#page-8-0) used specific labeling by fluorophores at the C-terminus of subunit ϵ and the N-terminus of subunit γ (Cy5 and Cy3, respectively; see Fig. 1) to monitor the conformational transitions in real time by measuring the fluorescence energy resonance transfer (FRET) from Cy3 to Cy5. They confirmed that ATP induces the transition of ε into the contracted state. They also found that when ATP was hydrolyzed to ADP by hexokinase, the reverse conformational change to the extended state of subunit ε occurred, in line with the suggested role of the ATP/ADP ratio. However, it remained ambiguous whether ADP exerts any direct effect on subunit ε , or whether it is the absence of ATP that matters. It was also unclear how the binding of ATP to subunit ε influences its conformational transitions.

We applied the FRET technique described above to investigate the conformational transitions of subunit ε in whole Bacillus PS3 F_0F_1 in native uncoupled membranes at 59 $^{\circ}$ C in the presence of both ATP and ADP.

MATERIALS AND METHODS

Expression and purification of F_1

Escherichia coli cells expressing Bacillus PS3 F_0F_1 (His-tagged at β -subunits, lacking subunit ε and with a cysteine introduced in position 3 of subunit γ , named $\gamma_{\text{Cys}}F_1-\Delta\varepsilon$) were grown and collected as described previously ([42,49\)](#page-8-0). DK8 E. coli cells (bglR, thi-1, rel-1, HfrPO1, Δ (uncBuncC), ilv:Tn10) lacking the wild-type (WT) F_0F_1 operon were used for expression. The $\gamma_{\text{Cys}}F_1-\Delta\varepsilon$ complex was purified by Ni-NTA affinity chromatography as described in the [Supporting Material](#page-7-0).

Bacillus PS3 ATP synthase subunit ε with a cysteine introduced in position 134 (ε_{Cys}) was expressed and purified as described previously [\(44,48\)](#page-8-0). Mutant ϵ EARA subunit ϵ was engineered from ϵ_{Cys} by substitution of residues ε Glu-83 and ε Arg-92 to alanines as described by Kato-Yamada and Yoshida ([44\)](#page-8-0).

Labeling of subunits γ

The purified $F_1\Delta\varepsilon$ with mutation S3C in subunit γ ($\gamma_{\text{Cys}}F_1-\Delta\varepsilon$) was treated for 1 h with 2 mM dithiothreitol (DTT) at 42°C. The sample was then passed through a NAP-10 gel filtration column equilibrated with buffer containing 100 mM KCl, 20 mM K_2HPO_4 , 2.5 mM EDTA, pH 6.6. Then the sample was diluted to 4 mg/mL protein by the same buffer, and Cy3-maleimide was added to a final concentration of 100 μ M from 5 mM stock solution in N,N-dimethyl formamide. After a 2 h incubation in darkness at room temperature, the reaction was stopped by addition of DTT to 2 mM.

Reconstitution of $F_1\Delta \varepsilon$ with subunit ε

E. coli cells expressing Bacillus PS3 ε_{Cys} (4 g, wet weight) were suspended in 70 mL of buffer containing 10% glycerol, 20 mM HEPES-KOH (pH 7.5), and 2.5 mM $MgCl₂$, passed twice through a French press, and mixed with 3 mL of $F_1\Delta \varepsilon$ labeled by Cy3-maleimide. The mixture was heated to 57°C for 10 min, and centrifuged at 190,000 \times g at 4°C for 1 h. The supernatant was applied to a NiNTA column, and reconstituted F_1 (with Cy3-labeled subunit γ and unlabeled subunit ε) was purified as described above. The purified protein was labeled by Cy5-maleimide according to the Cy3-maleimide labeling protocol.

The resulting F_1 labeled by Cy3 at γ Cys3, and by Cy5 at ε Cys-134 was concentrated as described above, and the excess Cy5 was removed on a NAP-10 column equilibrated with buffer containing 10% glycerol, 20 mM HEPES-KOH (pH 7.5), and 2.5 mM MgCl₂. The labeling efficiency

was checked by means of the absorption spectrum as described previously ([48\)](#page-8-0) and was found to be 50–60% for subunit γ and 95% for subunit ε . After purification was completed, the protein was concentrated to

2 mg/mL and stored at -80° C.

Reconstitution of F_1 into EDTA-treated membranes

Inverted *Bacillus* PS3 membrane vesicles stripped of F_1 were prepared by sonication of *Bacillus* PS3 cells in buffer containing 2 mM EDTA and 2 mM sodium succinate (see the [Supporting Material](#page-7-0) for details). Residual ATPase activity was inhibited by 7-chloro-4-nitrobenzo-2-oxa-1,3-diazole (NBD-Cl) treatment as described previously ([50\)](#page-8-0). For reconstitution, stripped membranes were mixed with an excess of purified F_1 , heated for 5 min to 60°C, and incubated for 30 min at room temperature. The excess F_1 was washed away by the same buffer in two sequential centrifugations $(190,000 \times g, 45 \text{ min}, 4^{\circ}\text{C})$. The reconstituted membranes were frozen in liquid nitrogen and stored at -80° C until they were used.

FRET measurements

FRET from Cy3 on subunit γ to Cy5 on subunit ε was monitored via Cy3 fluorescence on an FP-6500 fluorometer (Jasco, Tokyo, Japan) as described previously ([48\)](#page-8-0), with minor modifications (see the [Supporting Material\)](#page-7-0). Measurements were obtained in buffer containing 100 mM KCl, 1 mM MgCl2, 20 mM MES, 20 mM MOPS, 20 mM HEPES, 20 mM Tricine, 10% v/v glycerol, pH 7.9. The F_1 concentration in the cuvette was $3-10$ μ g/mL. In the case of reconstituted membranes, the total protein concentration was 20–60 μ g/mL, corresponding to F₁ concentrations in the range of 0.6–3.5 μ g/mL, as estimated by the initial fluorescence level. All FRET measurements were done at 59°C.

Measurement of ATPase activity

ATPase activity was measured via Phenol Red absorption changes as described previously ([51\)](#page-8-0), with minor modifications. See details in the [Supporting Material](#page-7-0).

RESULTS

ADP cannot reverse the ATP-induced conformational transition of ε

In previous studies of Bacillus PS3 enzyme, it was determined that 1), ATP induces the conformational transition of subunit ε from an extended to a contracted state; and 2), FIGURE 2 Effect of ATP and ADP on the conformational state of subunit ε , as monitored via fluorescence of Cy3 at 566 nm. Measurements were done at 59°C in buffer containing 100 mM KCl, 1 mM $MgCl₂$, 20 mM MES, 20 mM MOPS, 20 mM HEPES, 20 mM Tricine, 10% v/v glycerol, pH 7.9. Arrows indicate addition of nucleotides. Note the different timescales in panels A and B. Red traces: ATP added to 250μ M at time 0, no other additions. Purple traces: ATP added to $250 \mu M$ at time 0, ADP added later to 1 mM. Blue traces: ADP added to 1 mM at time 0, no other additions. Cyan traces: ADP added to 1 mM at time 0, ATP added later to 250 μ M. (A) isolated F_1 . (B) F_1 reconstituted with *Bacillus* PS3 inverted EDTA-treated membranes.

if all ATP is hydrolyzed to ADP, a reverse transition to an extended conformation occurs [\(42,48\)](#page-8-0). However, it was unclear which factor (i.e., the ATP/ADP ratio or just the ATP concentration) determines the conformation of ε . To clarify this issue, we investigated the effects of ATP and ADP in isolated F_1 and in F_0F_1 in inverted *Bacillus* PS3 membranes.

As can be seen in Fig. 2, the addition of ATP to 250 μ M caused a marked increase in the Cy3 fluorescence both in isolated F_1 and in F_0F_1 incorporated in *Bacillus* PS3 inverted membranes (red trace). It should be noted that the addition of MgATP complex was necessary to induce the transition. As documented in [Fig. S6,](#page-7-0) Mg-free ATP (as well as Mg^{2+} without ATP) induced no significant changes in Cy3 fluorescence.

When ADP alone was added to the sample to 1 mM (blue and cyan traces), no major change in the Cy3 fluorescence occurred. The slow, minor rise in the fluorescence observed after ADP addition was due to contaminating ATP in the commercial ADP used; this rise could be significantly decreased by the addition of a glucose-hexokinase trap (not documented).

When ADP was added to 1 mM after 250 μ M ATP (purple traces), no significant fluorescence decrease was observed. This result demonstrates that ADP failed to cause a reverse transition of ϵ from the ATP-induced contracted state.

The presence of 1 mM ADP also did not prevent subunit ε transition induced by 250 μ M ATP (compare *cyan* and *red* traces in Fig. 2). The extent of the fluorescence change was not diminished in the presence of ADP. The effects described were observed both for isolated F_1 (Fig. 2 A) and for F_0F_1 in *Bacillus* PS3 membranes (Fig. 2 B). The results imply that a >400-fold change in the ATP/ADP ratio had no major effect on the final conformation of subunit ε .

However, ADP lowered the rate of ATP-induced transition. The half-rise time for the fluorescence change increased from \approx 3 s (no ADP) to \approx 9 s (1 mM ADP) in isolated F₁, whereas in F_0F_1 the half-rise time increased from ≈ 15 to \approx 74 s (Fig. 2).

How much ATP is needed to induce the conformational transition of ϵ in F₁ and F_OF₁?

As documented in [Fig. 2,](#page-2-0) the addition of ATP caused an increase in Cy3 fluorescence in both isolated F_1 and the whole F_0F_1 . The extent of this increase was dependent on the final concentration of ATP added (Fig. 3). The dependence was different for isolated F_1 and the whole F_0F_1 in uncoupled membrane. The apparent K_d for ATP was 30 \pm 2 μ M in isolated F₁, but it increased to 141 \pm 8 μ M after reconstitution of F_1 with EDTA-treated *Bacillus* PS3 inverted membranes (Fig. 3). The Hill numbers, however, were similar: 1.60 \pm 0.14 in isolated F₁ and 1.76 \pm 0.13 in the whole F_0F_1 .

In full agreement with previous results ([48\)](#page-8-0), in isolated F_1 the kinetics of ATP-induced Cy3 fluorescence increase was apparently biexponential. The same was true for the whole F_0F_1 (Table 1; see also [Fig. S2\)](#page-7-0). Of note, the relative input of the faster and slower apparent components ($\approx 60\%$ and \approx 40%, respectively) was the same for isolated F₁ and the whole F_0F_1 within experimental error.

However, the time constant of each kinetic component was larger in the case of the whole F_0F_1 . The results in Table 1 indicate that binding of F_1 to F_0 slowed down the conformational transition of ε induced by 1 mM ATP ~6-fold.

FIGURE 3 ATP titration of subunit ε conformational transition as monitored via fluorescence of Cy3 at 566 nm in F_1 (*open circles*) and F_0F_1 (*solid* $circles$). Measurements were done at 59 \degree C in buffer containing 100 mM KCl, $1 \text{ mM } MgCl₂$, $20 \text{ mM } MES$, $20 \text{ mM } MOPS$, $20 \text{ mM } HEPES$, 20 mM Tricine, 10% v/v glycerol, pH 7.9. The fluorescence change was induced by addition of ATP to the respective final concentration $(x \text{ axis})$, and the change extent was measured after the plateau was reached (y axis, normalized by the fluorescence change extent after addition of ATP to 1 mM). Trace with water added instead of ATP was used as baseline. The inset shows an example of experimental traces in which ATP was added to a WT F_1 sample at time 0 to final concentrations of 1 mM, 250 μ M, 62.5 μ M, 31.25 μ M, 15.6 μ M, and 7.8 μ M (top to bottom). Each data point was measured in triplicate or more, and the standard deviation is plotted as error bars. Solid lines represent the best fits of the data points with a Hill function, dotted lines are fits with a Hill number of 2, and dashed lines are fits with a Hill number of 1.

TABLE 1 Kinetic parameters (time constants and relative inputs of components) of subunit ε conformational transitions induced by addition of ATP to 1 mM at 59° C

Sample	τ_1 , S	A1, $%$	τ_2 , S	A2, $%$	\boldsymbol{n}
WT F1	1.3 ± 0.3	$58 + 8$	9.6 ± 2.4	42 ± 8	9
WT F ₀ F ₁	7.8 ± 2.6	$61 + 4$	57 ± 22	$39 + 4$	-6
β D390A F ₁	1.2 ± 0.4	$71 + 8$	8.0 ± 2.9	$29 + 8$	7
β D390A F _o F ₁	3.0 ± 1.4	$64 + 6$	$18 + 4$	$36 + 6$	$\overline{4}$
β AALSAAA F ₁	0.6 ± 0.3	84 ± 4	5.7 ± 1.6	16 ± 4	7
β AALSAAA F _o F ₁	0.9 ± 0.2	$80 + 2$	17 ± 3	$20 + 2$	3
ϵ EARA F _o F ₁	1.8 ± 0.8	60 ± 2	28 ± 2	40 ± 2	$\overline{4}$

Traces were fitted with a biexponential function; n is the number of traces analyzed.

Role of the β DELSDED region in the conformational transition of ε

The electrostatic interactions of the negatively charged residues in the β DELSDED region of *Bacillus* PS3 ATP synthase were previously demonstrated to play an important role in the inhibition of ATP hydrolysis by subunit ε ([43\)](#page-8-0). In this work, we attempted to clarify their role in the conformational transitions of ε . We analyzed two mutants: one with a single substitution β D390A (first Asp in β DELSDED), and one with a full substitution β AALSAAA. According to the previous study (43) (43) , the β D390A mutation only slightly reduced (by 20%) the ATPase activity of the $\alpha_3\beta_3\gamma$ -complex, and diminished the inhibitory effect of subunit ϵ threefold. In the β AALSAAA mutant, the inhibitory effect of ε was reduced ninefold, but the ATPase activity of the $\alpha_3\beta_3\gamma$ -complex was sevenfold lower than in the WT.

As can be seen in Fig. $4A$ and B , both mutations resulted in acceleration of the ε transition induced by 1 mM ATP. The effect was clear in both isolated F_1 and the whole F_0F_1 . A detailed analysis revealed that the kinetics of transition in the mutants was also apparently biexponential for both isolated F_1 and the whole F_0F_1 (Table 1).

In the β D390A mutant, the slight acceleration observed in isolated F_1 was not due to the change in the time constants of transition kinetic components; rather, it reflected the increase in the fast component relative input (from $\approx 60\%$ in the WT to \approx 70% in the β D390A mutant). In contrast, the same mutation in the whole F_0F_1 led to an ~3-fold decrease in both time constants, leaving the relative inputs unaffected.

In the fully substituted β AALSAAA mutant, the transition time constants for isolated F_1 were ~2-fold smaller than in the WT, and the relative input of the fast component increased from $\approx 60\%$ to $\approx 80\%$ (Table 1). The strongest difference was observed in F_0F_1 for the fast component time constant, which dropped more than eightfold (from 7.8 s in the WT to 0.9 s in the β AALSAAA mutant). The slow component was also accelerated (57 s vs. 17 s).

The dependence of the ATP-induced transition extent on ATP concentration in the β D390A and β AALSAAA mutants is shown in [Fig. 5.](#page-4-0) In the case of the β D390A mutant ([Fig. 5](#page-4-0) A), the dependence was very similar to that of the

FIGURE 4 Effect of mutations on the kinetics of subunit ε conformational transition induced by 1 mM ATP in F_1 (A and C) and F_0F_1 (B and D). Measurements were done at 59°C in buffer containing 100 mM KCl, 1 mM MgCl2, 20 mM MES, 20 mM MOPS, 20 mM HEPES, 20 mM Tricine, 10% v/v glycerol, pH 7.9. (A and B) β DELSDED mutants (red trace: WT; green trace: β D390A mutant; blue trace: β AALSAAA mutant). (C and D) ε EARA mutant (red trace: WT; cyan trace: ε EARA mutant). The inset in panel C shows the same traces during first 20 s after addition of ATP. ATP added at time 0 to a final concentration of 1 mM. Traces were scaled to the same maximal extent. Note the different timescales.

WT [\(Fig. 3\)](#page-3-0) for both isolated F_1 (*open circles*) and F_0F_1 (solid circles). In the case of the β AALSAAA mutant (Fig. 5 B), the apparent K_d was markedly lower than in the WT for both the isolated F₁ (5.6 μ M vs. 30 μ M) and F_OF₁ (15 μ M vs. 141 μ M). The Hill number in mutant F_OF₁ was also significantly decreased: a value of $n \approx 0.8$ was observed (compared to $n \approx 1.8$ in the WT).

Role of ATP binding directly to subunit ϵ in its conformational transitions

To investigate the role of ATP binding directly to subunit ε , we substituted two residues involved in this binding $(\varepsilon \text{Glu-})$ 83 and ϵ Arg-92 [\(47](#page-8-0)); *orange spheres* in [Fig. 1](#page-1-0) A) to alanines. The resultant mutant was named ϵ EARA. Isolated subunit ϵ with this double mutation had very low ATP binding; the apparent K_d was >1 mM at 25°C (F. Kadoya and Y. Kato-Yamada, unpublished observations). Taking into account the fact that the WT showed >100 -fold lower ATP binding at 60° C than at 25° C ([48\)](#page-8-0), there must have been negligible, if any, ATP binding to the ϵ EARA mutant at 59 $\mathrm{^{\circ}C}$ with 1 mM ATP in this study.

The ATPase activity of isolated ϵ EARA F₁ (1 mM ATP, 59°C) was indistinguishable from that of the WT F_1 . However, in the whole ϵ EARA F_OF₁ (isolated or incorporated in liposomes), the rate of ATP hydrolysis was 60% \pm 7% of the WT rate.

FIGURE 5 ATP titration of subunit ε conformational transition as monitored via fluorescence of Cy3 at 566 nm in F_1 (*open circles*) and F_0F_1 (*solid* circles). Measurements were done as in [Fig. 3](#page-3-0). Solid lines represent the best fits of the data points with a Hill function, dotted lines are fits with a Hill number of 2, and dashed lines are fits with a Hill number of 1. (A) β D390A mutant. (B) β AALSAAA mutant. (C) ϵ EARA mutant. Standard deviation is plotted as error bars for data points measured in triplicate; other data points are the average of two measurements. Solid lines represent the best fits of the data points with a Hill function, dotted lines are fits with a Hill number of 2, and dashed lines are fits with a Hill number of 1.

A comparison of the ATP-induced conformational change of ε in the WT and in the mutant enzyme is shown in Fig. 4, C and D. In the isolated F_1 , the mutation slowed

TABLE 2 Apparent K_d and Hill number for the dependence of the donor fluorescence change extent on ATP concentration

Sample	Apparent K_d , μ M	Hill number
$WT F_1$	30 ± 2	1.60 ± 0.14
WT F ₀ F ₁	141 ± 8	1.76 ± 0.13
β D390A F ₁	$30 + 2$	1.60 ± 0.13
β D390A F _o F ₁	124 ± 9	1.65 ± 0.17
β AALSAAA F ₁	5.6 ± 0.8	1.57 ± 0.33
β AALSAAA F _O F ₁	15 ± 2	0.81 ± 0.19
ϵ EARA F ₁	31 ± 2	1.39 ± 0.11
ϵ EARA F _O F ₁	136 ± 19	1.24 ± 0.24

the transition down. As can be seen in [Fig. 4](#page-4-0) C , the kinetics was no longer biexponential. Immediately after the addition of ATP, a fast, unresolved change in the fluorescence was observed (see the inset in [Fig. 4](#page-4-0) C). This fast component accounted for $\approx 20\%$ of the total fluorescence change. It was followed by a biexponential increase of the signal with time constants of \approx 20 s and \approx 320 s.

In the whole F_0F_1 , the mutation effect was the opposite of that observed in isolated F_1 : the conformational transition of ϵ was faster in ϵ EARA. No unresolved initial phase was observed, and the kinetics could be fitted with a biexponential function ([Table 1](#page-3-0)).

The ATP titration curves for the ϵ EARA mutant are shown in [Fig. 5](#page-4-0) C. The apparent K_d values were very close to these of the WT F_1 (Table 2). The Hill number was somewhat reduced in the mutant, especially in the whole F_0F_1 ($n \approx$ 1.2 vs. $n \approx 1.8$ in the WT).

Influence of the subunit ϵ conformation on the ATPase activity of F_0F_1 in Bacillus PS3 membranes

It has been established that the extended conformation of subunit ε is inhibitory and the contracted hairpin conformation is not [\(42](#page-8-0)). Moreover, in our previous study on isolated F_1 in the presence of the ATP regenerating system, it was determined that the fast kinetic component of the Cy3 fluorescence change correlated with the increase in the ATPase activity of the enzyme [\(48](#page-8-0)). However, we believed that additional experiments in membrane samples at 59°C were necessary.

To investigate the effect of the subunit ε conformation on F_0F_1 ATPase activity, *Bacillus* PS3 membranes stripped of F_1 with EDTA and treated with NBD-Cl to block any residual ATPase activity were reconstituted with four types of F₁: 1), the WT γ Cy3\eCy5-labeled F₁ used throughout this work; 2), the $\varepsilon^{\Delta C}$ -F₁ with the C-terminal domain of subunit ε deleted (used as a control with zero inhibitory effect of ε); 3), the unlabeled $\gamma_c \varepsilon_c$ -F₁ treated with CuCl₂ in the absence of ATP to fix subunit ε in the extended state by a cross-link as described previously ([42\)](#page-8-0) (used as a control with a maximal inhibitory effect of ε); and 4), the latter $\gamma_c \varepsilon_c$ -F₁ cross-linked enzyme treated with DTT to break the cross-link (used as a control mimicking the WT enzyme).

FIGURE 6 ATPase activity of F_0F_1 in *Bacillus* PS3 membranes and its correlation with the Cy3 fluorescence changes. Measurements were done at 59° C. See details in text. (A) Dependence of the initial ATPase activity on ATP concentration. Bacillus PS3 membranes reconstituted with different F_1 samples were diluted to 9 μ g protein/mL with buffer containing 1 mM tricine-KOH (pH 8.05), 150 mM KCl, 1 mM $MgCl₂$, 50 μ M phenol red. ATP hydrolysis was started by addition of ATP to the desired final concentration (x axis, note the log scale). Open triangles: $\varepsilon^{\Delta C}$ F_OF₁; black circles: γ Cy3/ ϵ Cy5 F_OF₁; black squares: $\gamma_c \epsilon_c$ F_OF₁ with subunit ϵ fixed in the extended state by a cross-link; gray squares: $\gamma_c \varepsilon_c$ F_OF₁ without cross-link. The inset shows the same data (low ATP concentration part) with higher resolution. Each point was measured at least in triplicate. (B) Relative effect of subunit ε C-terminus on ATPase activity at different ATP concentrations. Gray columns: The activity of $\gamma Cy3/\varepsilon Cy5$ F_OF₁ was divided by the activity of $\epsilon^{\Delta C}$ F_OF₁ and normalized to one at an ATP concentration of 1 mM. Solid circles: ATP titration of subunit ε conformational transition in γ Cy3/ ε Cy5 F_0F_1 monitored via fluorescence of Cy3 at 566 nm (same conditions as for ATP hydrolysis measurements). Solid line: Hill function best fit of $F_OF₁$ ATP titration data duplicated from [Fig. 3](#page-3-0).

After reconstitution with F_1 , the initial ATPase activity was measured in each sample at different ATP concentrations (Fig. 6 A). In all samples, the ATPase activity decreased when the ATP concentration was lowered. However, this decrease was less pronounced in $\varepsilon^{\Delta C}$ -F_OF₁ than in other samples. We normalized the ATPase activity by the value at 1 mM ATP and plotted the ratio of normalized activity in the γ Cy3\ ϵ Cy5-labeled F_OF₁ to normalized activity in

the $\varepsilon^{\Delta C}$ sample. The relative decrease in ATPase activity at lower ATP concentrations was more than twofold larger in γ Cy3\cCy5-labeled F_OF₁ than in ε ^{AC}-F_OF₁ [\(Fig. 6](#page-5-0) B, gray columns). To directly compare this effect with the ATPinduced FRET change, we also measured the FRET change in γ Cy3\sCy5-labeled F_OF₁ sample in the same buffer that was used for ATPase activity measurements ([Fig. 6](#page-5-0) B, solid circles). The results indicated a good correlation between the relative FRET change and the relative ATPase activity.

DISCUSSION

We extended the FRET technique that was successfully applied for real-time monitoring of subunit ε conformational transitions in isolated F_1 to the whole F_0F_1 integrated in Bacillus PS3 inverted membrane vesicles. The enzyme was studied in a native lipid environment at 59°C (close to 65° C, the optimal growth temperature for *Bacillus* PS3). No ATP regenerating system or glucose\hexokinase trap was added during the measurements, so both ATP and ADP were present in the medium, as they would be in vivo. Unfortunately, the reconstituted membranes were not well coupled at 59°C (see the [Supporting Material\)](#page-7-0). Therefore, the results describe F_0F_1 incorporated into uncoupled membrane with no significant proton-motive force present.

The results shown in [Fig. 2](#page-2-0) indicate that in the absence of ATP, subunit ε was in the extended state (highest FRET efficiency, lowest Cy3 fluorescence), irrespective of the presence of ADP. ATP induced the transition of ε to a contracted state, again irrespective of the presence of ADP (although ADP slowed the transition down \sim 5-fold; [Fig. 2](#page-2-0) B). Moreover, the addition of ADP in fourfold excess did not reverse the ATP-induced transition. These findings suggest that the equilibrium between the contracted and extended conformations of ε was influenced only by the ATP concentration, and not by ATP/ADP ratio.

The ATP titration of the subunit ε conformational transi-tion ([Fig. 3](#page-3-0)) revealed that the apparent K_d for transition is 141 μ M. ATP titration of the ATPase activity in the γ Cy3\ ε Cy5-labeled F_OF₁ used for FRET measurements, and in $\varepsilon^{\Delta C}$ -F_OF₁ that lacked the inhibitory C-terminal domain of ε ([Fig. 6\)](#page-5-0) indicated that the observed conformational transition correlated well with the increase in the subunit ε inhibitory effect. Unless some unknown factors significantly alter the behavior of subunit ϵ in vivo, these results imply that the hairpin noninhibitory conformation of ϵ prevails under normal physiological conditions (when intracellular ATP concentration is in the millimolar range). In contrast, the extended inhibitory state of ε is expected to occur only when the intracellular ATP concentration decreases to $\leq 100-200$ μ M. It is tempting to speculate that in Bacillus PS3 ATP synthase, subunit ε serves as an ''emergency break'' that strongly inhibits the ATP hydrolysis activity of the enzyme when the intracellular ATP pool is almost exhausted. Further experiments are necessary to prove or dismiss this hypothesis.

A Hill number higher than one ([Fig. 3](#page-3-0) and [Table 2\)](#page-5-0) indicated that binding of at least two ATP molecules per enzyme was necessary to induce the transition of ε . A noninteger Hill number suggested that the mechanism of transition induction was complex and could not be described in the framework of a simple two-site cooperative binding model. From a physiological point of view, a higher Hill number results in a narrower interval of ATP concentrations where the transition occurs, i.e., an increase in the Hill number shifts the response of the system from a gradual to an ''all-or-nothing'' reaction triggered by a decrease/increase of ATP concentration below/above a certain limit.

The list of known nucleotide-binding sites on F_0F_1 includes three catalytic sites and three noncatalytic sites on the $\alpha_3\beta_3$ hexamer, and one site on subunit ϵ . However, we note that 1), in isolated F_1 the deletion of the noncatalytic sites did not abolish the ATP-induced transitions of subunit ε [\(34,48\)](#page-8-0); 2), as can be seen in [Figs. 4 and 5,](#page-4-0) the deletion of the nucleotide-binding site on subunit ε did not abolish these transitions either; and 3), previous studies revealed that in isolated F_1 the deletion of the catalytic sites resulted in the loss of an ATP-induced conformational change of subunit ϵ , as detected by cross-linking [\(47](#page-8-0)). Therefore, it is likely that MgATP binding to the catalytic sites is the primary event that induces the conformational transition of subunit ε , as was proposed previously ([48\)](#page-8-0).

Taking into consideration that the apparent K_d value was 141 μ M for the ϵ transition ([Table 2](#page-5-0)), it is probable that all three catalytic sites have to be occupied to drive the subunit ϵ transition from the inhibitory extended state. It should be noted that in the β AALSAAA mutant F_0F_1 , the apparent K_d value was lower than that of the WT by one order of magnitude, and there was no indication of positive cooperativity in ATP binding (Hill number < 1 ; see [Fig. 5](#page-4-0) B, solid circles, and [Table 2\)](#page-5-0). Probably in this case, binding of ATP to one catalytic site (or two) was sufficient to induce the transition of ε .

The apparent K_d value of 15 μ M for the β AALSAAA mutant from the ATP titration experiment ([Table 2](#page-5-0)) suggests that interactions of acid residues in β DELSDED are involved in stabilizing the extended conformation of subunit ε , as was hypothesized in an earlier study [\(52](#page-8-0)). It should be noted that the first Asp in β DELSDED does not seem to be involved in such interactions, because its replacement to Ala had no detectable effect on the ATP titration curve in either F_0F_1 or isolated F_1 ([Table 2\)](#page-5-0). However, this replacement diminishes the inhibitory effect of ε by a factor of 3 [\(43](#page-8-0)). Our results also indicate that this replacement accelerated the ATP-induced transition of subunit ε ([Table 1\)](#page-3-0). It is therefore likely that the interactions between the first Asp in β DELSDED and the C-terminus of ε occurred in some intermediate, transitional state that differed from the stable, fully extended conformation, although they were also inhibitory.

The kinetics of the ATP-induced conformational change of subunit ε was apparently biexponential in both F_1 and F_0F_1 [\(Table 1](#page-3-0) and Fig. S2). Such kinetics could be expected if the transition of the ε C-terminus were a complex process with two or more stages. Therefore, it is probable that there are some intermediate conformation(s) between the fully extended state, in which the C-terminus of ε is in close proximity to the N-terminus of γ , and a fully contracted hairpinlike state. Of note, the relative inputs of the fast and slow kinetic components were the same for isolated F_1 and F_0F_1 ([Table 1\)](#page-3-0). Therefore, it seems likely that the conformational transition of subunit ε follows the same pattern and has the same intermediate state(s) in isolated F_1 and whole F_0F_1 .

In this work, we were also interested in clarifying the role of ATP binding to ε . A previous study revealed that alanine replacements of residues involved in ATP binding to ϵ enhanced the inhibitory effect ([47\)](#page-8-0). In agreement with these results, ATPase activity in ϵ EARA mutant F_0F_1 was also diminished (60% of the WT activity). This was in apparent contradiction to the results documented in [Figs. 4 and 5](#page-4-0). On the one hand, we observed that in the ϵ EARA mutant, ATP induced a conformational transition of ε similar to that in the WT enzyme. Therefore, one could expect full inhibition relief at millimolar ATP concentrations. On the other hand, in ϵ EARA F_OF₁, a 40% inhibition of the steady-state ATP hydrolysis was observed at 1 mM ATP (i.e., at this ATP concentration, the mutant subunit ε was no longer in the extended state, but was still inhibitory). Our results suggest that the transition of ϵ from the extended state to some contracted state was necessary but was not sufficient to eliminate the inhibition of ATPase activity. A specific hairpin-like fold stabilized by bound ATP was probably essential for full activity. Unlike subunit ε from E. coli F_0F_1 , *Bacillus* PS3 subunit ε requires ATP binding to keep its C-terminus in a hairpin fold (46) (46) . Thus, in the ϵ EARA mutant, the subunit ε C-terminus probably remained unfolded and inhibitory even at high ATP concentrations (state II in [Fig. 1](#page-1-0) B). Such behavior is indistinguishable from the transition to the hairpin conformation (state III in [Fig. 1](#page-1-0) B) revealed by the FRET measurements, because in both cases the distance from Cy5 at the C-terminus of ε to Cy3 at the N-terminus of subunit γ significantly increased.

Our results and earlier data can be explained by the following simple scheme (Fig. $1 B$): At low ATP concentrations, ε is in the inhibitory extended state (state I). Binding of the MgATP complex to the catalytic sites induces conformational changes that drive the C-terminal domain of ε out of the $\alpha_3\beta_3$ hexamer ([Fig. 1](#page-1-0) B, $I\rightarrow II$). In the hypothetical intermediate state (state *II*), the C-terminus of ε may be able to adopt different conformations, some of which could be inhibitory of ATPase activity (as was observed in ϵ EARA F_0F_1). Other, noninhibitory conformations include the contracted hairpin state. In the WT enzyme (but not in the ϵ EARA mutant), the noninhibitory hairpin state is stabilized by binding of ATP ([Fig. 1](#page-1-0) B, $II \rightarrow III$). If the ATP concentration later drops below the apparent K_d of ATP binding to ϵ $(\approx 200 \ \mu M$ at 59°C [\(48](#page-8-0))), ATP will dissociate from subunit ϵ , thereby destabilizing the hairpin fold [\(Fig. 1](#page-1-0) B, III \rightarrow II). A further decrease in ATP concentration below the apparent K_d of the ε conformational transition (141 μ M at 59°C; see [Table 2\)](#page-5-0) leads to a reverse transition of subunit ϵ [\(Fig. 1](#page-1-0) B, $II\rightarrow I$) and a strong inhibition of ATP hydrolysis.

SUPPORTING MATERIAL

Additional materials and methods, results and discussion, six figures, and references are available at [http://www.biophysj.org/biophysj/supplemental/](http://www.biophysj.org/biophysj/supplemental/S0006-3495(09)01630-0) [S0006-3495\(09\)01630-0.](http://www.biophysj.org/biophysj/supplemental/S0006-3495(09)01630-0)

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